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► To cite this version:

Katia Abbaci, Allel Hadjali, Ludovic Liétard, Daniel Rocacher. A Similarity Skyline Approach for Handling Graph Queries - A Preliminary Report. International Workshop on Graph Data Management Techniques and Applications (GDM'11), in Conjunction with The IEEE International Conference on Data Engineering (ICDE), Apr 2011, Germany. pp.112-117. hal-00670641

HAL Id: hal-00670641

<https://hal.science/hal-00670641>

Submitted on 15 Feb 2012

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A Similarity Skyline Approach for Handling Graph Queries - A Preliminary Report

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Abstract—One of the fundamental problems in graph databases is similarity search for graphs of interest. Existing approaches dealing with this problem rely on a single similarity measure between graph structures. In this paper, we suggest an alternative approach allowing for searching similar graphs to a graph query where similarity between graphs is rather modeled by a vector of scalars than a unique scalar. To this end, we introduce the notion of *similarity skyline* of a graph query defined by the subset of graphs of the target database that are the *most similar* to the query in a *Pareto sense*. The idea is to achieve a d -dimensional comparison between graphs in terms of d local distance (or similarity) measures and to retrieve those graphs that are maximally similar in the sense of the Pareto dominance relation. A diversity-based method for refining the retrieval result is proposed as well.

I. INTRODUCTION

Graphs have become increasingly important in modeling complex structured data in many recent real applications. These applications include Bioinformatics [1], [2], Pattern Recognition [3], XML documents [4], Chemical compounds [5], Social networks [6], etc. All these applications indicate the importance and the broad usage of graph databases. One can broadly classify queries against graph databases into two categories [7]: (1) *Graph containment search* and (2) *Graph similarity search*. The former consists of the following two sub-problems: (i) *subgraph containment search*: given a graph database $D = \{g_1, g_2, \dots, g_n\}$ and a graph query q , retrieve all graphs $g_i \in D$ such that q is a subgraph of g_i (i.e., $q \subseteq g_i$); (ii) *supergraph containment search*: given a graph database $D = \{g_1, g_2, \dots, g_n\}$ and a graph query q , retrieve all graph $g_i \in D$ such that q is a supergraph of g_i (i.e., $q \supseteq g_i$). Both sub-problems consider the procedure of checking *subgraph isomorphism*, known to be NP-Complete. Many query processing approaches using indexing techniques have been developed to reduce the search space and then efficiently solve these two sub-problems [8], [9], [10], [11].

As for the second category (i.e., graph similarity search), which consists in retrieving all the graphs of the database that are structurally similar to a given graph query, has emerged

as new trend due to the following reasons [12], [13]. Firstly, many real graph datasets are noisy and incomplete in nature, so approximate, rather than exact, *graph matching* is required. Secondly, many graph applications prefer approximate matching results rather than exact ones as they can provide more information such as what might be missing or spurious in a query or in a graph database. A number of approaches therefore have been proposed to support similarity queries on graph databases, see [14], [15] and [12] in the case of subgraph queries and [16] in the case of supergraph queries. The common point of all those approaches is the fact that they impose a single measure to evaluate graph similarity. However, a graph is a complex structure by nature and involves various basic features. It is then difficult to give a meaningful definition of graph similarity using only a single index.

In this paper, we advocate that for graph similarity to be efficiently assessed, several indices are required. Each index is dedicated to measure a local distance (or similarity) between two graphs pertaining to one aspect in the graph structure. Therefore, graph similarity is now characterized by a vector of local distance measures (where each measure expresses a *feature similarity*) instead of a single measure. By this way, one can preserve information about similarity on several features when comparing two graphs.

We propose an approach based on the notion of similarity skyline to support graph similarity search. Roughly speaking, the *similarity skyline* of a graph query is defined by the subset of graphs of the target database that are the *most similar* to the query in a *Pareto sense*. The idea is to achieve a d -dimensional comparison between graphs in terms of d local distance (or similarity) measures and to retrieve those graphs that are *maximally similar* in the sense of a defined *similarity-dominance relation*. In summary, we made the following contributions in this paper:

- We introduce the notion of graph compound similarity and then define the *similarity-dominance relationship* between graphs.
- Based on that relationship, we give a formal definition

of the *graph similarity skyline*, i.e., graphs of the target database that are maximally similar to a graph query in a Pareto sense.

- To reduce the resulting skyline (which is often quite large), we propose a method to extract a subset which is *as diverse as possible*, but with an acceptable size.

The rest of the paper is organized as follows. Section 2 provides some preliminary notions. Related work is discussed in Section 3. Section 4 describes some well-known measures for graph similarity and their semantic properties. In Section 5, we introduce the notion of similarity skyline to support graph similarity queries. Section 6 proposes a detailed example. Section 7 presents a method for refining the retrieval result. Section 8 concludes the paper.

II. PRELIMINARIES

A. Reminder About Skyline Queries

Skyline queries [17] are a popular and powerful paradigm for extracting interesting objects from a multi-dimensional dataset. They rely on Pareto dominance principle which can be defined as follows:

Definition 1. Let r be a set of d -dimensional data points and $p = (p_1, p_2, \dots, p_d)$ and $q = (q_1, q_2, \dots, q_d)$ two points of r . p is said to dominate (in the Pareto sense) q iff on every dimension $p_i \leq q_i$ (for $1 \leq i \leq d$) and on at least one dimension $p_j < q_j$.

For simplicity and without loss of generality, we assume that the smaller the value p_i , the better. We say then that p *dominates* (is preferred to) q and we denote this by $p \succ q$.

Definition 2. The skyline of r is the set of points which are not dominated by any other point.

Skyline queries compute the set of Pareto-optimal tuples in a relation, i.e., those tuples that are not dominated by any other tuple in the same relation.

Example 1. Consider a database containing information about hotels as shown in Table I (where dimension $d = 2$).

TABLE I
SAMPLE HOTELS

Hotel	Price(€)	Distance (Km)
H_1	4.0	150
H_2	3.0	110
H_3	2.5	240
H_4	2.0	180
H_5	1.7	270
H_6	1.0	195
H_7	1.2	210

Consider a person who looks for a hotel that is as close as possible to the beach and having a low price. One can check that the resulting skyline S contains H_2 , H_4 and H_6 . For instance, H_1 is dominated by H_2 , and H_7 by H_6 .

B. Some Basic Definitions

Definition 3 (Graph). A graph g is defined as a 4-tuple (V, E, L, l) where V is the set of vertices, E is the set of edges, L

is the set of labels and l is a labeling function that maps each vertex or edge to a label in L .

For ease of presentation, graphs refer here to undirected labeled graphs. Note that different nodes could have the same label and the size of g is defined as $|g| = |E(g)|$ (i.e., the size of a graph is the number of its edges).

Definition 4 (Graph isomorphism). Given two graphs $g = (V, E, L, l)$ and $g' = (V', E', L', l')$, g is *isomorphic* to g' (denoted by $g \approx g'$) if there exists a *bijection* $f : V \rightarrow V'$, such that

- 1) $\forall v \in V, f(v) \in V'$ and $l(v) = l'(f(v))$ and;
- 2) $\forall (u, v) \in E, (f(u), f(v)) \in E'$, and $l(u, v) = l'(f(u), f(v))$.

Definition 5 (Subgraph isomorphism). Given two graphs $g = (V, E, L, l)$ and $g' = (V', E', L', l')$, g is *subgraph isomorphic* to g' if there exists an *injection* $f : V \rightarrow V'$ such that

- 1) $\forall v \in V, f(v) \in V'$ and $l(v) = l'(f(v))$ and;
- 2) $\forall (u, v) \in E, (f(u), f(v)) \in E'$ and $l(u, v) = l'(f(u), f(v))$.

Definition 6 (Subgraph v.s. supergraph). Given two graphs $g = (V, E, L, l)$ and $g' = (V', E', L', l')$, g is called a *subgraph* of g' (or g' is a *supergraph* of g), denoted as $g \subseteq g'$ (or $g' \supseteq g$), if there exists a subgraph isomorphism from g to g' .

Definition 7 (Maximum common subgraph, mcs). Given two graphs g_1 and g_2 , the *maximum common subgraph* of g_1 and g_2 is the largest (i.e., the maximum number of selected vertices) connected subgraph of g_1 that is subgraph isomorphic to g_2 , denoted as $g' = mcs(g_1, g_2)$.

III. RELATED WORK

Our proposal can be related to the works in the areas of skyline queries and similarity queries on graph databases.

Skyline queries. They have received a lot of attention over the recent years. Several research efforts have been made to develop efficient algorithms and to introduce different variants for skyline queries [18], [19], [20], [21]. Up to our knowledge, no work related to skyline queries exists in a graph data context, except the recent work by Zou et al. [22] where dynamic skyline queries in a large graph have been studied. In our case, a different kind of skyline (i.e., similarity skyline) over a set of graphs (rather than a single large graph) is investigated.

Similarity queries. Similarity search of graphs is a vital operation in many recent applications. As indicated in Section I, this kind of graph search is conducted thanks to similarity queries that aim at finding graphs in the target database that are similar, but not necessarily isomorphic, to a given graph query. A number of approaches have been developed to support similarity queries. *Grafil* [14] performs substructure similarity search in a large scale graph database. It returns all the graphs of the database that approximately contain the graph query. *C-Tree* [15] is another tool for subgraph similarity search which focuses on the edit distance between the query and its candidate matches. *Tale* [12], unlike most previous graph matching tools which treat every node in a

graph equally, proposes an innovative matching technique that distinguishes nodes by their importance in the graph structure. This technique first matches the important nodes of a graph query, and then progressively extends these matches. Recently, Shang *et al.* [16] have proposed a technique to deal with supergraph queries where the notion of maximum common subgraph plays a key role. The problem of interest is converted into a σ -missing subgraph detection problem, where σ is the error tolerance threshold. All the graphs of the queried database such that the mcs-based distance measure to the graph query considered is less than σ , are returned as answers. Both *C-Tree* and *Tale* rely on the *edit distance* to measure similarity between graphs whereas the works done in [14] and [16] use the notion of *maximum common subgraph* for computing that similarity.

As can be seen all the approaches that support similarity queries on graph data make use of a unique index to measure similarity between two graphs. So doing, similarity between two graph structures is not wholly captured since some similarities pertaining to some features of graph are missed. This is mainly due to the fact that each index of graph similarity can be seen as a local measure that expresses only a resemblance w.r.t. one aspect in a graph structure (see Section IV). Compared with the above work, our approach, on the one hand, relies on a compound similarity measure between graphs and, on the other hand, returns a set of similarity dominant graphs in a Pareto sense to answer a graph query.

IV. GRAPH SIMILARITY MEASURES: SOME SEMANTIC PROPERTIES

Several models have been proposed [23], [24], [25] to measure the similarity (or distance) between two graphs. Hereafter, we present the most widely accepted measures to determine similarities between graphs¹.

A. Graph Edit Distance

Graph edit distance [15], [23] is based on graph edit operations needed to transform one graph to another. Generally, the set of edit operations considered includes: insertion or deletion of a vertex/edge and relabeling of a vertex/edge. Each edit operation is associated with a cost (a non-negative real number) according to the amount of distortion that it introduces in the transformation. Let e_op be an edit operation and $c(e_op)$ its cost. The cost of a sequence of edit operations, $s = (e_op_1, \dots, e_op_n)$ is given by

$$c(s) = \sum_{i=1}^n c(e_op_i).$$

The choice of elementary edit operations and their cost represent a difficult task in practice. The cost of a transformation of an element to another can be regarded as a distance function between the two elements. We assume here a *uniform* distance measure: the distance between two vertices/edges is 1 if they have different labels; otherwise it is 0.

¹Due to space limitation, the computational complexity analysis of each measure is not addressed here.

Definition 8 (Graph edit distance). The edit distance between two graphs g_1 and g_2 is equal to the *minimum cost*, taken over all sequences of edit operations, that transform g_1 into g_2 , i.e.,

$$Dist_{Ed}(g_1, g_2) = \min_{s \in E_op} c(s) \quad (1)$$

where E_op denotes the set of all sequences of edit operations that transform g_1 into g_2 .

The smaller $Dist_{Ed}(g_1, g_2)$, the more similar the two graphs. One can easily check that the edit distance between *isomorphic* graphs is zero.

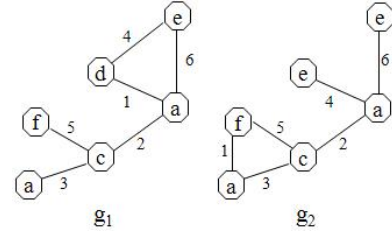


Fig. 1. Examples of labeled graph

Example 2. Let us consider the labeled graphs of Fig.1. The sequence of edit operations those are necessary for transforming g_1 into g_2 is: (i) one edge deletion, (ii) one edge relabeling, (iii) one vertex relabeling, (iv) one edge insertion. By considering uniform distance measures, one can check that this sequence is the best one (i.e., the shortest). So, $Dist_{Ed}(g_1, g_2) = 4$.

It is worth noticing that, in a graph database querying context, this distance measure provides information on features that both a graph of the target database and the graph query at hand disagree.

B. Mcs-Based Distance

Bunke *et al.* [24] have developed another kind of measure for graph similarity. It is based on the *maximum common subgraph* (mcs).

Definition 9 (Similarity based on the mcs). Given two graphs g_1 and g_2 , the graph similarity based on the mcs is defined as,

$$Sim_{Mcs}(g_1, g_2) = \frac{|mcs(g_1, g_2)|}{\max(|g_1|, |g_2|)},$$

where $|mcs(g_1, g_2)|$ denotes the number of edges in $mcs(g_1, g_2)$.

Clearly, the larger the *mcs* of two graphs the greater their similarity. The measure Sim_{Mcs} is normalized (i.e., $0 \leq Sim_{Mcs}(g_1, g_2) \leq 1$) since $|mcs(g_1, g_2)| \leq \max(|g_1|, |g_2|)$. Now, the graph distance measure, $Dist_{Mcs}$, derived from Sim_{Mcs} writes:

$$Dist_{Mcs}(g_1, g_2) = 1 - Sim_{Mcs}(g_1, g_2) \quad (2)$$

Such a measure is proved to be a metric in [24] and leads to a distance in $[0, 1]$.

The major advantage of the mcs-based approach is the fact that it does not require the use of any cost function, thereby avoiding the main drawback of edit-distance-based approach.

Example 3. Let us come back to Example 2. The mcs -based distance measure between g_1 and g_2 is calculated as follows. First, the $mcs(g_1, g_2)$ is identified, see Fig. 2. Then, by applying (2), we obtain

$$Dist_{Mcs}(g_1, g_2) = 1 - \frac{|mcs(g_1, g_2)|}{\max(|g_1|, |g_2|)} = 0.33,$$

where $|mcs(g_1, g_2)| = 4$ and $\max(|g_1|, |g_2|) = 6$.

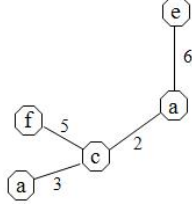


Fig. 2. Mcs of g_1 and g_2

From a database querying point of view, this kind of similarity conveys information on the amount of features that both a graph of the queried database and a graph query agree.

C. Gu-Based Distance

Graph union (Gu)-based distance measure, introduced by Wallis *et al.* [25], is based on the idea of graph union. Graph union (rather than the larger of two graphs) is used to model the size of the problem.

Definition 10 (Gu-based similarity). Given two graphs g_1 and g_2 , the graph similarity based on graph union is defined as,

$$Sim_{Gu}(g_1, g_2) = \frac{|mcs(g_1, g_2)|}{|g_1| + |g_2| - |mcs(g_1, g_2)|},$$

where the denominator represents the size of union of the two graphs in the set theoretic sense².

This similarity measure is also normalized and its behaviour is somewhat close to Sim_{Mcs} . It is easy to see that $Sim_{Gu}(g_1, g_2) \leq Sim_{Mcs}(g_1, g_2)$ holds as well (which means that Sim_{Gu} is a stronger measure than Sim_{Mcs}). The use of graph union [25] is motivated by the fact that changes in the size of the smallest graph that keep $mcs(g_1, g_2)$ constant are not taken into account in $Sim_{Mcs}(g_1, g_2)$ whereas the measure $Sim_{Gu}(g_1, g_2)$ does take this variation into account.

The graph distance measure derived from Sim_{Gu} can be written as:

$$Dist_{Gu}(g_1, g_2) = 1 - Sim_{Gu}(g_1, g_2) \quad (3)$$

It was also proved to be a metric and gives values in $[0, 1]$.

Example 4. Let us again consider the graphs provided in Example 2. Using (3), the Gu-based distance measure between g_1 and g_2 is

$$Dist_{Gu}(g_1, g_2) = 1 - \frac{|mcs(g_1, g_2)|}{|g_1| + |g_2| - |mcs(g_1, g_2)|} = 0.50,$$

where $|mcs(g_1, g_2)| = 4$ (see Example 3) and $|g_1| = |g_2| = 6$.

In a database querying context, this type of similarity gives also information about the number of agreements between a graph of the queried database and a graph query.

²This similarity measure looks like the Jaccard index used to measure similarity between two sets A and B , i.e., $J(A, B) = |A \cap B| / |A \cup B|$.

V. GRAPH SIMILARITY SKYLINE

This section is devoted to define the notion of *similarity skyline* for supporting graph similarity search without the need for specifying a global similarity measure between graph structures.

From now on, we assume that graph similarity is a compound notion, i.e., in order to assess similarity between graphs we consider a vector of distance measures. Each measure can be regarded as a local similarity expressing the extent to which two graphs are similar w.r.t. some features or aspects.

Definition 11 (Graph Compound Similarity, GCS). Let g and g' be two graphs, a graph compound similarity between g and g' is a vector of local distance measures denoted by

$$GCS(g, g') = (Dist_1(g, g'), Dist_2(g, g'), \dots, Dist_d(g, g')),$$

where $Dist_i(g, g')$, for $i = 1, \dots, d$, stands for a local graph distance measure.

Let now $D = \{g_1, g_2, \dots, g_n\}$ be a graph database and q a graph similarity query (i.e., this means that the user is interested in graphs of D that are the most similar to q). Since a global similarity between graphs is not available, the idea is to proceed with a d -dimensional comparison between graphs in terms of d (local) distance measures to retrieve graphs that are maximally similar in the sense of the following similarity-dominance relation.

Definition 12 (Similarity-dominance relation). Given a graph query q and two graphs g and g' , we say that g' is *similarity-dominated* by g in the context of q , denoted by $g \succ_q g'$, iff the following two statements hold:

- 1) $\forall i \in \{1, \dots, d\}, Dist_i(g, q) \leq Dist_i(g', q)$,
- 2) $\exists k \in \{1, \dots, d\}, Dist_k(g, q) < Dist_k(g', q)$.

Roughly speaking, the relation $g \succ_q g'$ holds if g is not less similar to q than g' in all dimensions and (strictly) more similar to q than g' in at least one dimension. One can observe that g is potentially more interesting than g' as a retrieval graph. Therefore, the set of graphs that are the most similar to q are those that are not dominated (in the sense of Definition 12). Such graphs, called *Pareto-optimal graphs*, represent what we denote by the *graph similarity skyline* (GSS):

$$GSS(D, q) = \{g \in D \mid \nexists g' \in D, g' \succ_q g\} \quad (4)$$

where $g' \succ_q g$ means that g is similarity-dominated by g' .

To illustrate the above approach, we provide in the next section an example where $d = 3$. $GCS(g, q)$ is then a vector of three components expressed by the local distance measures described in Section IV, i.e.,

$$GCS(g, q) = (Dist_{Ed}(g, q), Dist_{Mcs}(g, q), Dist_{Gu}(g, q)).$$

VI. AN ILLUSTRATIVE EXAMPLE

Let $D = \{g_1, g_2, g_3, g_4, g_5, g_6, g_7\}$ be a graph database and q a graph similarity query, as shown in Fig. 3. In order to provide the most interesting answers to q , one can compute the graph similarity skyline $GSS(D, q)$. It is easy to see that $|g_1| = 6$, $|g_2| = 7$, $|g_3| = 7$, $|g_4| = 6$, $|g_5| = 8$, $|g_6| = 9$, $|g_7| =$

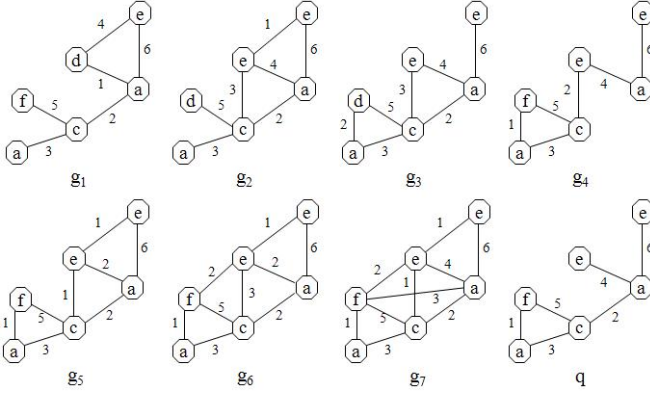


Fig. 3. The graph database D and the graph query q

10 and $|q| = 6$. Table 2 summarizes the values of $|mcs(g_i, q)|$, for $i = 1, \dots, 7$.

TABLE II
INFORMATION ABOUT $|Mcs(g_i, q)|$

	$ Mcs(g_i, q) $
(g_1, q)	4
(g_2, q)	4
(g_3, q)	4
(g_4, q)	3
(g_5, q)	5
(g_6, q)	5
(g_7, q)	6

TABLE III
DISTANCE MEASURES

	$Dist_{Ed}(g_i, q)$	$Dist_{Mcs}(g_i, q)$	$Dist_{Gu}(g_i, q)$
(g_1, q)	4	0.33	0.50
(g_2, q)	4	0.43	0.56
(g_3, q)	3	0.43	0.56
(g_4, q)	2	0.50	0.67
(g_5, q)	3	0.38	0.44
(g_6, q)	4	0.44	0.50
(g_7, q)	4	0.40	0.40

Now the graph similarity vectors $GCS(g_i, q)$, for $i = 1, \dots, 7$, are shown in Table III. By applying (4), the set of Pareto optimal graphs, i.e. the graph similarity skyline, is given by $GSS(D, q) = \{g_1, g_4, g_5, g_7\}$.

One can easily check that g_2 (resp. g_3) $\notin GSS(D, q)$ since it is dominated by g_7 (resp. g_5) and $g_6 \notin GSS(D, q)$ since it is dominated by g_1 . Thus, the graphs of D that are maximally similar to q are g_1, g_4, g_5 and g_7 . Indeed,

- Graph g_1 is the most interesting w.r.t. the measure $Dist_{Mcs}$. This is due to the two following reasons: i) g_1 satisfies a maximum number of features required by q than other graphs with the same size; ii) g_1 and q are of the same size. But, g_1 is the less interesting w.r.t. to superfluous and missing features.

- Graph g_4 is the best w.r.t. the measure $Dist_{Ed}$. This means that it is the most interesting w.r.t. to the numbers of disagreements with q . On the other hand, g_4 is much less satisfactory w.r.t. to the agreements with q in the sense of the mcs notion.
- Graph g_7 is the most interesting w.r.t. the measure $Dist_{Gu}$. This is due to the fact that $g_7 \supset q$. But, it is the less interesting w.r.t. a superfluous feature-based criterion.
- Graph g_5 may be a good compromise between the three measures $Dist_{Ed}$, $Dist_{Mcs}$ and $Dist_{Gu}$.

Let us now take a look at the results obtained when using only a single similarity measure between graphs. If we are interested in the best $k (= 3)$ answers, g_3 is then returned for instance by the edit-distance-based approach as answer to the user, but with the skyline-based approach g_3 is not returned as answer since g_5 does better than it.

VII. REFINING GRAPH SIMILARITY SKYLINE

One of the problems that may arise when computing the set GSS (and a skyline in general) is its size which is often quite large. From a user point of view, it is very desirable to have a suitable criterion to select a small interesting subset of graphs of the skyline GSS. One solution is to use the criterion of diversity [26] to select a subset of graphs which is *as diverse as possible* and then provide the user with a picture of the whole set GSS.

Let S be a subset of GSS. The diversity of S means that the graphs it includes should be dissimilar amongst each other. The goal is to extract from GSS a subset \mathbb{S} of size k (a user-defined parameter) with a *maximal diversity*. Adapted from [27], the proposed approach defines the diversity of S ($\subseteq GSS$) of size k by a vector $Div(S) = (v_1, v_2, v_3)$ such as

$$v_i = \min\{Dist_i(g, g') | g, g' \in S\},$$

where $Dist_1 = Dist_{N-Ed}$ (the normalized version of the distance $Dist_{Ed}$ obtained using the function $f(x) = x/(1+x)$), $Dist_2 = Dist_{Mcs}$ and $Dist_3 = Dist_{Gu}$. The value v_i expresses the diversity in the i^{th} dimension of the subset S .

To identify the subset \mathbb{S} of interest, we consider all subsets $S \subseteq GSS$ with $|S| = k$ (i.e., the size of S is k) as candidates and apply the following steps:

Step 1. For each dimension i ($i = 1, \dots, 3$), rank-order all candidates subsets S in decreasing way according to their diversity v_i in that dimension. Let $rank_i(S)$ be the rank of S w.r.t. i^{th} dimension. Rank value 1 means the best diversity value and rank value M means the worst diversity value (M is the number of subsets of size k of the set GSS).

Step 2. Evaluate a candidate S by:

$$val(S) = \sum_{i=1, \dots, 3} rank_i(S).$$

The subset that minimizes this criterion (i.e., minimizes the sum of its positions in all ranks) is considered as a subset with a maximal diversity. So, \mathbb{S} is characterized by

$$val(\mathbb{S}) = \min_S val(S)$$

where $S \subseteq GSS$ and $|S| = k$.

Example 5. Let us come back to the example given in section VI, where $GSS(D, q) = \{g_1, g_4, g_5, g_7\}$. Assume now that the user is interested in the best k ($= 2$) graphs w.r.t. the diversity criterion. One can easily check that the set of all candidates contains 6 subsets of size k , see Table IV.

TABLE IV
CANDIDATES WITH THEIR DIVERSITY

	v_1	v_2	v_3
$S_1 = \{g_1, g_4\}$	0.86	0.67	0.80
$S_2 = \{g_1, g_5\}$	0.83	0.50	0.60
$S_3 = \{g_1, g_7\}$	0.87	0.60	0.67
$S_4 = \{g_4, g_5\}$	0.80	0.62	0.73
$S_5 = \{g_4, g_7\}$	0.83	0.70	0.77
$S_6 = \{g_5, g_7\}$	0.75	0.50	0.61

Now, steps 1 and 2 lead to the results depicted in Table V.

TABLE V
EVALUATION OF ALL CANDIDATES.

(a) Ranks ($r_i = rank_i$).				(b) $Val(S_i)$.	
	r_1	r_2	r_3		$\sum_{i=1,\dots,3} r_i$
S_1	2	2	1	S_1	5
S_2	3	5	6	S_2	14
S_3	1	4	4	S_3	9
S_4	4	3	3	S_4	10
S_5	3	1	2	S_5	6
S_6	5	5	5	S_6	15

From Table V-(b), one can easily see that $val(S_1)$ is the minimal value. So, $\mathbb{S} = S_1 = \{g_1, g_4\}$.

VIII. CONCLUSION

In this paper, we have proposed an alternative approach to support graph similarity search. The key concept of this approach is the notion of graph similarity skyline we introduced. This kind of skyline allows retrieving all graphs of the queried database that are not dominated in the sense of the similarity-dominance relation defined. Namely, graphs those are maximally-similar to the graph query at hand. Each answer graph is provided to the user with a vector of scores showing different similarities pertaining to different features. We have also shown how to select a maximally diverse subset of a graph similarity skyline.

We plan to conduct some experiments on real-life data to demonstrate the effectiveness and efficiency of the approach. To this end, a system implementing it is underway.

ACKNOWLEDGMENTS

This work was supported in part by the National Agency for Research under project AOC on the reference ANR-08-CORD-009, and Brittany region.

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